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C; Keywords: transcription regulation
                                                                                                                                                     R:Park J.S.; Luethy, J.D.; Wang, M.G.; Fargnoli, J.; Fornace Jr., A.J.; McBride, O.W.; Gene 116, 259-267, 1992
A;Title: Tsolation, Characterization and chromosomal localization of the human GADD153 A;Reference number: JC1169; MUID:92339899
A;Accession: JC1169
                                                                                                                 A; Molecule type: mRNA
A; Residues: 1-168 < PAR>
                                                                                                                                                                                                                                                                               DNA-damage-inducible protein GADD153 - human
C;Species: Homo sapiens (man)
C;Date: 05-Mar-1993 #sequence_revisido 05-Mar-1993 #text_change 21-Jul-2000
                                      A; Gene: GADD153
                                                        C; Genetics
                                                                          C; Comment: This protein is
                                                                                                 A; Cross-references:
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A; Note: TLS is a synonym fo
C; Keywords: fusion protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           A:Crbss-references: EMBL:X71427; NID:g395919; PIDN:CAA50558.1; PID:g4210364 C:Comment: This sequence is the chimeric product of a translocation mutation C:Genetics:
A:Gene: GADD153/FUS
                                                                                                                                                                                                                                                                    C:Accession: JC1169
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            A; Molecule type: mRNA
A; Residues: 1-462 < RAB>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       A;Title: Fusion of the dominant negative transcription regulator CHOP with a novel A;Reference number: S36157; MUID:93350637 A;Accession: S36158
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     A;Cross-references: GB:S62138; NID:g386258; PIDN:AAB27103.1; PID:g386159 R;Rabbitts, T.H.; Forster, A.; Larson, R.; Nathan, P. Nature Genet. 4, 175-180, 1993
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A; Molecule type: mRNA
A; Residues: 1-462 <CRO>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        A:Title: Fusion of CHOP to a novel RNA-binding protein in human myxoid liposarcoma. A:Reference number: $33798; MUID:93288139
A:Accession: $33798
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            R;Crozat, A.; Aman, P.; Mandahl, N., Ron, D. Nature 363, 640-644, 1993
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     C; Date: 22-Nov-1993 #sequence_revision 38 Nov-1995 #: C; Accession: S33798; S36158
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                FUS/CHOP mutant fusion protein - human N:Alternate names: TLS/CHOP mutant fusion protein
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Best Local S
Matches 85
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                                                                                                                                                                                                                                                                                                                                                                                                                                       354 AWLTEEEPEPAEVTSTSQSPHSPDSSQSSLAQEEEEEDQG 393
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         294 MAAESLPFSFGTLSSWELEAWYEDLQEVLSSDENGGTYVSPPGNEEEESKIFTTLDPASL 353
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Conservative
                                                                 GB:S40706; NID:g252001; PIDN:AAB22646.1; tein is responsible for the growth arrest
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          84.7%;
85.0%;
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Pred. No. 1.6e-33;
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                                                                                                                                                                                                                                                                                                                                                          1873, PAD 40
                                                                                        PID: g252002
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Matches
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A; Title: Sequence and analysis of chromosome 2 of the plant Arabidopsis thaliana. A; Reference number: A84420; MUID:20083487
A; Accession: F84770
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             A;Molecule type: mRNA
A;Residues: 1-237 <IZS/
A;Cross-references: DDBJ:D63707
C;Comment: This protein translocates to the nucleus and directly functions in mitogen
F;1-98/Region: hath #status predicted
F;155-170/Region: nuclear location signal
                                                                                            A; Map position:
                                                                                                                                                                                             A; Molecule type: DNA
A; Residues: 1-263 <STO>
                                                                                                                                                                                                                                                                                                                                             R;Lin, X.; Kaul, S.; Rounsley, S.D.; Shea, T.P.; Benito, M.I.; Town, C.D.; Fujii, C.Y M.; Koo, H.; Woffat, K.S.; Cronin, L.A.; Shen, M.; Vanaken, S.E.; Umayam, L.; Tallon, euss, D.; Nierman, W.C.; White, O.; Elsen, J.A.; Salzberg, S.L.; Fraser, C.M.; Venter
                                                                                                                                                                                                                                                                                                                                                                                                                                           hypothetical protein At2g35600 [Imported] - Arabidopsis thaliana C;Species: Arabidopsis thaliana (mouse-ear cress) C;Date: 02-Feb-2001 #sequence_revision 02-Feb-2001 #text_change 02-Feb-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     RESULT
F84770
                                                                                                                            A; Gene: At2g35600
                                                                                                                                                C; Genetics:
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JC5660
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C;Date: 17-Nov-1997 #sequence_revision 17-Nov-1997 #text_change 17-Mar-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                R; Izumoto, Y.; Kuroda, T.; Harada, H.; Kishimoto, T.; Nakamura, H. Biochem. Biophys. Res. Commun. 238, 26-32, 1997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           R; Izumoto,
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                                                                                                                                                                                                                                                                                                                                                                                                                                      ;Accession: F84770
                             Query Match
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         187 -- GERPLPVEVEKNS-TPSEPDSGQGPPAEEEEGEEE 220
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                                                                                                                                                            GB:AE002093; NID:g3608128; PIDN:AAC36161.1; GSPDB:GN00139
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85.0%;
18.5%; Score 94; [
25.8%; Pred. No. 0.
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Pred. No. 0.053;
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Pred. No. 3.5e-33;
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                 DB 2; Length 263;
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Local Similarity

31;

Conservative

19;

Mismatches

Indels

28;

Gaps

.12; 42;

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Title:
Perfect score:
Sequence:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      OM protein - protein search, using sw model
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Maximum Match 100%
Listing first 45 summaries
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Maximum DB seq length: 2000000000
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                                  Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                        PIR_71:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:*
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Gapop 10.0 , Gapext 0.5
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508
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Copyright (c) 1993 - 2000 Compugen Ltd.
SUMMARIES
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29	28	27	26	25	24	23	22	21	20	19	18	17	16	. 15	14	13	12	11	10	9	89	7	σ	u	4	ω	2	<b></b> -	No.	Result
73.5	73.5	73.5	73.5	74	74	75	75.5	76	77.5	77.5	78.5	80	80.5	80.5	80.5	80.5	81	81.5	81.5		ω	83.5	94	97	421	430.5	435	508	Score	
	14.5	14.5						15.0	15.3	15.3	15.5		15.8	15.8	15.8	15.8	15.9	16.0	16.0			16.4	18.5	9	2		85.6	100.0	Match	Query
550	382	382	187	418	344	310	3375	296	1840	872	440	240	1755	1054	440	303	297	1624	214	1755	440	440	263	237	168	462	168	168	Length	
ν	N	2	N	N	Ν	N	N	Ν	N	N	N	N	N	N	N	ν	ν	2	Ν	2	N	N	N	N	N	4	N	N	DB.	
T29919	T08243	JQ1122	<b>В86704</b>	T15827	T40167	146987	T19821	A54527	T30250	T25186	S52895	A55055	869969	D96519	S69970	A86443	S76306	T25592	T39559	S45736	S45737	S49765	F84770	JC5660	JC1169	S33798	A34096	S26148	ID	
hypothetical pr	gas-vesicle operon	gas-vesicle prote	_	_	_	0			- 17	hypothetical pr	- 1	riv	TyB protein - y	g		0	hypothetical pr	hypothetical pr	٠.	protein -	protein -	- 1	hypothetical pr	hepatoma-derived	DNA-damage-inducib	FUS/CHOP mutant	DNA damage inducib	nuclear protein	Description	
prote	ron	tei	prote	prote	prote	in	prote	Pla	mous	prote	yeas	g	yeas	tei	yeas	int	prote	prote	5	yeas	yeas	yeas	prote	g	di	fu	dic	C)	ì	

C; Keywords: transcription regulation

Query Match 85.6%; Score 435; DB 2; Length 168; Best Local Similarity 88.7%; Pred. No. 1.6e-34; Matches 86; Conservative 3; Mismatches 8; Indels

0; Gaps

0;

) 1990 M.C.; id by	Db 1 MANESLEFILE VSSMELEAW INDIGSTRONG TO THE INTERPRETATION OF TH		RESULT 1  \$26148  nuclear protein chop-10 - mouse  C; Species: Mus musculus (house mouse) C; Date: 13-Jan-1995 #sequence_revision 13-Jan-1995 #text_chan C; Accession: \$26148 R; Ron, D.; Habener, J.F. Genes Dev. 6, 439-453, 1992 A; Title: CHOP, a novel developmentally regulated nuclear prot A; Reference number: \$26148; MUID:92192456 A; Accession: \$26148 A; Status: preliminary A; Molecule type: mRNA A; Residues: 1-168 < RON> A; Cross-references: EMBL: X67083; NID:g50406; PIDN:CAA47465.1; C; Keywords: transcription regulation	ALIGNMENTS	30 73.5 14.5 1114 2 150222 31 73 14.4 113 2 C89828 32 73 14.4 648 2 H42687 33 72.5 14.3 349 2 T42665 34 72 14.2 617 2 A56051 35 72 14.2 1171 2 T13065 36 72 14.2 2232 2 T34434 37 71.5 14.1 2210 2 139410 38 71.5 14.1 1210 2 139410 39 71.5 14.1 1211 2 A568198 40 71 14.0 309 2 T00503 41 71 14.0 309 2 T00503 42 71 14.0 318 2 C70550 44 70.5 13.9 185 2 C70550 44 70.5 13.9 555 2 A56560 45 70.5 13.9 672 2 T12524
#text_change 05-Nov-1999; Luethy, J.D.; Papathanasiou, M.; growth arrest signals and DNA-dama	SPGNEEEESKTFTTLDPASL 60 M	2; Length 168; 41; 0; Indels 0; Gaps 0;	<pre>#text_change 05-Nov-1999 uclear protein that dimerizes with CaA47465.1; PID:g50407</pre>		deltaEFI - chicken conserved hypothet probable wD-40 rep glycoprotein - ate myocyte nuclear fa pip82 protein - fr hypothetical prote chromogranin A pre AF-4 protein, spli serine/proline-ric probable MYB famil probable transcrip delta-crystallin/E hypothetical protein hypothetical hypothetical hypothetical hypothetical hypothetical hypothetical

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OM protein - protein search, using sw model

June 18, 2002, 15:09:51; Search time 11.83 Seconds (without alignments) 324.027 Million cell updates/sec

Title: Perfect score:

Sequence: US-09-637-550-3 508 1 MAAESLPFTLETVSSWELEA......RSPDSSQSSMAQEEEEEEQG 99

Scoring table:

BLOSUM62 Gapop 10.0 , Gapext 0.5

105224 seqs, 38719550 residues

Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database : SwissProt\_40:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

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Query Match 100.0%; Score 508; DB 1; Length 168; Best Local Similarity 100.0%; Pred. No. 1.4e-41; Matches 99; Conservative 0; Mismatches 0; Indels 0; Gaps

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## ALIGNMENTS

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DDIT3 OR CHOP OR GADD153.
Cricetulus longicaudatus (Long-tailed hamster) (Chinese hamster).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         01-APR-1990 (Rel. 14, Created)
01-APR-1990 (Rel. 14, Last sequence update)
01-NOV-1997 (Rel. 35, Last annotation update)
Growth arrest and DNA-damage-inducible protein GADD153 (DNA-damage
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation -
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA-damaging agents.";
Mol. Cell. Biol. 9:4196-4203(1989).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Fornace A.J. Jr., Nebert D.W., Hollander M.C., Luethy J.D., Papathanasiou M., Fargnoli J., Holbrook N.J.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TISSUE=Ovary
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     the European Bioinformatics Institute. There are no restrictions use by non-profit institutions as long as its content is in modified and this statement is not removed. Usage by and for contents
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                -1- FUNCTION: INHIBITS THE DNA-BINDING ACTIVITY OF C/EBP AND LAP FORMING HETERODIMERS THAT CANNOT BIND DNA.
-1- FUNCTION: MAY PLAY A ROLE IN THE INHIBITION OF GROWTH AFTER
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 MEDLINE=90066424;
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GA15_HUMAN
                                                                                                                                                                                                                                                                                                                             InterPro; IPR001871; bZIP.

SMART; SM00338; BRLZ; 1.

Growth arrest; Nuclear protein; Transcription regulation; DOMAIN 92 95 POLY-GLU.
                                                                                                                                                                                                                                                                                                                                                                                                                                                       entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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                                                                                                               61 AWLTEEPGPTEVTRTSQSPRSPDSSQSSMAQEEEEEE 97
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   PRT;
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01-JUN-1994 (Rel. 29, Created)
01-JUN-1994 (Rel. 29, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
16-OCT-2001 (Rel. 40, Last annotation update)
Growth arrest and DNA-damage-inducible protein GADD153 (DNA-damage inducible protein)

Crowth arrest and DNA-damage-inducible protein)

(CHOP).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 This SWISS-PROT entry is copyright. It is produced through a collaboratic between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on it use by non-profit institutions as long as its content is in no was modified and this statement is not removed. Usage by and for commercia
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               MEDLINE=92339899; PubMed=1339368; Park J.S., Luethy J.D., Wang M.G., Fargnoli J., Fornace A.J. Jr., McBride O.W., Holbrook N.J.;
                                                                                                                                          CONFLICT
SEQUENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Rabbitts T.H., Forster A., Larson R., Nathan P.;
"Fusion of the dominant negative transcription regulator CHOP with a novel gene FUS by translocation t(12;16) mailgnant liposarcoma.";
Nat. Genet. 4:175-180(1993).
-i- FUNCTION: INHIBITS THE DNA-BINDING ACTIVITY OF C/EBP AND LAP BY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQUENCE FROM N.A. MEDLINE=93288139; PubMed=8510758;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                              Growth arrest; Nuclear protein; Transcription regulation; Repressor;
                                                                                                                                                                                                                                                                                                                                                                   MIM; 126337; -. MIM; 151900; -.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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DOMAIN 93 97 POLY-GLU.
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85.0%;
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695.069 Million cell updates/sec
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                                                                                                                                                                                                                                                                                                                                                                                                                              sp_human:*
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                                                                                                                                                                                                                                                                                                                                                                                                                                                sp_bacteria: *
                                                                                                                                                                                                                                                                                                       sp_unclassified:*
sp_rvirus:*
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sp_archeap:*
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    168
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082281
09F865
09F864
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1 0923W3
                      Q9XBQ7
Q9F861
Q9F866
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   Q03964
Q12266
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Q62804
Q62857
Q9CYA4
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09f865 enterococcu
09f864 enterococcu
09f863 enterococcu
09f863 rattus norv
0923w3 rattus norv
09x867 enterococcu
09f861 enterococcu
09f866 enterococcu
09f866 enterococcu
09f866 enterococcu
09f866 saccharomyc
012266 saccharomyc
                                                                                                      Q91yw9 mus musculu
Q62804 rattus norv
Q62857 rattus norv
Q92ya4 mus musculu
Q9jk87 mus musculu
Q82281 arabidopsis
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Q91YW9
AC Q9

Mus musculus (Mouse). Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus NCBI\_TaxID=10090;

O91YW9 PRELIMINARY; PRT; 168 AA. 091YW9; 01-DEC-2001 (TrEMBLrel. 19, Created) 01-DEC-2001 (TrEMBLrel. 19, Last sequence update) 01-DEC-2001 (TrEMBLrel. 19, Last annotation update) DNA-DAWAGE INDUCTBLE TRANSCRIPT 3.

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Q9xt15	09u7e8	015417	Q95jv7	Q9c0e4	Q9ngz2	Q962b3	09dg1	Q9etj5 c	091mg	06181	Q9xu67	09hch8	09v8b2	09qy79	012112	09:69	012391	69560	P74789	091rh	095q78	095a77	Q9dbd5	091ym9	042932	Q9f862	Q9h7h3	012490	
caenorhabdi	caenc	homo sa	macac	homo	homo sapien	elicoverpa	l fuğu rubrip	orynebacte	ratt	8 mus musculu		homo sapien	drosophila		accharomyc	8 arabidonsis			synechocy	2 raphanus sa	caenorhabdi	aen	Bull	9			homo sapie	saccharomvc	

## ALIGNMENTS

Db  Qy  Db  RESULT  RESULT  QC 2804  AC Q  DT Q  DT Q	RP RP RA RA SQ SQ SQ Ma
1 MAASSLPFTLETVSSWELEAWYEDLOEVLSSDENGGTYISSPGNEEEESKTFTTLDPASL 60 61 AWLITEEPGPTEVTRTSQSPRSPDSSQSSMAQEEEEEGG 99	SEQUENCE FROM N.A.  Strausberg R.; Strausberg R.; Strausberg R.; Submitted (SEP-2001) to the EMBL/GenBank/DDBJ databases. EMBL; BC013718; AAH13718.1; EMBL; BC013718; AAH13718.1; SEQUENCE 168 AA; 19190 MW; CC993B7957F43160 CRC64;  Ouery Match  98.68; Score 501; DB 11; Length 168; Best Local Similarity 99.0%; Pred. No. 3.le-44; Matches 98; Conservative 0; Mismatches 1; Indels 0; Gaps 0;  1 MARESLPFTLETYSSWELEAMYEDLQEVISSDEIGGTYISSPGNEEEESKTFTTLDPASL 60

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RESULT
Q9CYA4
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Best Local
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Best Local
                                                                                                                                                                                                                                                     STRAIN-SPRAGUE DAWLEY; TISSUE-BRAIN;
Jin K.L., Chen J., Simon R.P., Graham S.H.;
submitted (SEP-1995) to the EMBL/GenBank/DDBJ databases
EMBL; U36994; AAA87944.1; -.
InterPro; IPR001871; bZIP.
                                                                                                                                                                                                                                                                                                                                                                                                            01-NOV-1996 (TREMBLIEL 01, 01-NOV-1996 (TREMBLIEL 01, 01-JUN-2001 (TREMBLIEL 17,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Chen C.H., Nusscnzweig A., Li G., Ling C.C.; "Cloning of rat GADD153 and its expression in submitted (AUG-1995) to the EMBL/GenBank/DDBJ
                                                                                                                                                                                                                                                                                                                                                           Eukaryota; Metazoa; Chordata; Mammalia; Eutheria; Rodentia;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Q62857
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Submitted (JUN-1995) to the EMBL; U30186; AAA73629.1; -. InterPro; IPR001871; bZIP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Nussenzweig A.;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                NCBI_TaxID=10116;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Eukaryota; Metazoa;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Rattus norvegicus (Rat).
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  Q9CYA4
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                                                                                                                                                                                                                                                                                                                                               NCBI_TaxID=10116;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               61 AWLTEEPGPTEVTRTSQSPRSPDSSQSSMAQEEEEEEEQG 99
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                                                                                       AWLIEEPGPTEVTRTSQSPRSPDSSQSSMAQEEEEEEQG 99
                                                                                                                MAAESLPFAFETVSSWELEAWYEDLQEVLSSDEIGGTYISSPGNEEEESKTFTTLDPASL 60
                                                                                                                                        MAAESLPFTLETVSSWELEAWYEDLQEVLSSDEIGGTYISSPGNEEEESKTFTTLDPASL 60
                                                                 AWLTEEPGPAEVTSTSQSPRSPDSSQSSMAQEEEEEDQG
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    PRELIMINARY;
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                                                                                                                                                                                                                                  19013 MW;
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94.9%;
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94.9%;
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Last annotation update)
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                                                                                                                                                                           Score 485; UB 1.,
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Pred. No. 1.4e-42;
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                                                                                                                                                                                                                                                                                                                                                             Craniata; Vertebrata; Euteleosto
Sciurognathi; Muridae; Murinae;
    PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  2516D983D40EA6E1 CRC64;
                                                                                                                                                                                                                                    251716CD712BA6E1 CRC64;
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RESULT
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01-OCT-2000
01-OCT-2000
01-JUN-2001
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01-JUN-2001 (TrEMBLrel. 17,
01-JUN-2001 (TrEMBLrel. 17,
01-JUN-2001 (TrEMBLrel. 17,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MGD; MGI:1194494; Hdgf.
InterPro; IPR000313; PWWP.
Pfam; PF00855; PWWP; 1.
SMART; SM00293; PWWP; 1.
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STRAIN=C57BL/6J; T
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                                                                                                                                                                                   01-OCT-2000 (TrEMBLrel. 15, Created)
01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-UNN-2001 (TrEMBLrel. 17, Last annotation update)
TDRM1 PROTEIN (UNKNOWN) (PROTEIN FOR MGC:6452).
                                                                                                                                                                                                                                                                            Q9JК87
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EMBL; AK017863; BAB30979.1; -
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                 SEQUENCE FROM N.A. STRAIN=BALB/C; TIS Zhao Y., Chen W.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                SEQUENCE
                                                                                      Eukaryota; Metazoa; Chordata; Mammalia; Eutheria; Rodentia; NCBI_TaxID=10090;
                                                                                                                                                Mus musculus (Mouse).
                                                                                                                                                                     TURMI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            "Functional annotation of a full-length mouse cDNA collection.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Hayashizaki Y.;
                                                                                                                                                                                                                                                                                                                                                                       187 -- GERPLPVEVEKNS-TPSEPDSGQGPPAEEEEGEEE 220
                                                                                                                                                                                                                                                                                                                                                                                                                                              132 SSDEEGKLVIDEPAKEKNEKGTLKRRAGDVLEDSPKRPKESGDHEEEDKEIAALE----
                                                                                                                                                                                                                                                                                                                                                                                                           62 WLTEEPGPTEVTRTSQSPRSPDSSQSSMAQEEEEEEQ 98
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Local
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       30;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Similarity
of novel gene related to thymus development.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  237 AA;
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                                                                                                                                                                                                                                                                                PRELIMINARY;
                                      TISSUE=THYMUS;
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                   Wang
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30.9%;
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Last annotation update)
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Pred.
                                                                                                                               Craniata; Vertebrata; Euteleostomi;
                                                                                                                                                                                                                                                                                  PRT;
                                                                                                              Sciurognathi; Muridae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ------GNEEEESKTFTTLDPASLA 61
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  OAE1CF574DA1733C CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Mismatches
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Result
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Maximum Match 100%
Listing first 45 summaries
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Maximum DB seq length: 200000000
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Searched:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Scoring table:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Title:
Perfect score:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          OM protein - protein search, using sw model
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                                                                                                                                                                                                                                                                                                                                                                            Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                                                                                                            Score
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(without alignments)
365.447 Million cell updates/sec
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167
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AAW80745
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AAG32233
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                  Human secreted pro
Mouse liver cancer
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## ALIGNMENTS

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diagnostic, forensic, gene therapy and chromosome mapping procedures -	obtaining cDNAs and genomic DNAs that correspond to 5'ESTs and for	New nucleic acid that is a 5' expressed sequence that (5' FST) for		N-PSDB; AAC01625.	WPI; 2000-500381/45.		Dumas Milne Edwards J. Duclert A. Giordano J.		(GEST ) GENSET.	20" EED 1999			21-FEB-2000; 2000EP-0200610.		06-SEP-2000.		EP1033401-A2.	,	Homo sapiens.	and the state of t	dene therapy: chromosome manning	Human: 5' EST: expressed sequence that secreted protects of the total transfer	TOWNS STATE OF THE SEA TO NO. 3/00.	Liman corrected process, Growth was cross		06-OCT-2000 (first entry)		AAG01619;		AAG01619 standard; Protein; 80 AA.	AAG01619	

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   03-DEC-1998 (first entry)
This represents the mouse liver cancer cell-derived growth factor (mHDGF). HDGF facilitates nuclear transfer. The invention provides
                                                Claim 17; Pages 17-18; 22pp; Japanese
                                                                               New nucleus-transfer signal introducing human liver cancer cell-derived growth factor to nucleus - and new recombinant DNA, mutant and transformed E. Coli and animal cells
                                                                                                                                               WPI; 1998-535025/46.
N-PSDB; AAV55933.
                                                                                                                                                                                                                                                                    25-FEB-1997;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Nucleus-transfer signal peptide; HDGF-NLS; HDGF protein; mouse; human; liver cancer cell-derived growth factor; nuclear transfer.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Mouse liver cancer cell-derived growth factor (mHDGF).
                                                                                                                                                                                                 (SEKI ) SEKISUI CHEM IND CO LTD
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                                                                                                                                                                                                                                                                                                                                                                   /note= "the DNA sequence encoding the amino acids
    in this region are transposed"
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83.8%;
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Best Local
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23-MAR-1999

25-MAR-1999

26-AAR-1999

06-AAR-1999

06-AAR-1999

07-AAR-1999

21-AAR-1999

21-AAR-1999

23-AAR-1999

23-AAR-1999

23-AAR-1999

24-AAR-1999

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29-AA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           17-0CT-2000
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           termination sequence.
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                                                                                                                                                                                                                                                                990S-0130449.
990S-0130510.
990S-0130891.
990S-0131449.
990S-0132048.
990S-0132048.
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99US-0123548.
99US-0125788.
99US-0132863.
99US-0134256.
99US-0134219.
99US-0134219.
99US-0134221.
99US-0134370.
99US-0134768.
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99US-0132485.
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99US-0128234
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99US-0126785
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